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(54) Title: SPBRANCHED1A OF SOLANUM PENNELLII AND TOMATO PLANTS WITH REDUCED BRANCHING COMPRISING THIS HETEROLOGOUS SPBRANCHED1A GENE

(57) Abstract: The present invention relates to a new sequence of *BRANCHED1a* gene, the *SpBRANCHED1a* gene of *Solanum pennellii*, and to an expression product of said gene, to a vector, host cell, cell culture, recombinant expression system, or tomato plants comprising said *SpBRANCHED1a* gene. The present invention also refers to the heterologous use of the gene in tomato plants, to produce a plant having fewer and/or shorter axillary shoots than a control plant, and to a method for producing tomato plants having fewer and/or shorter axillary shoots than a control plant.



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SPBRANCHED1A OF SOLANUM PENNELLII AND TOMATO PLANTS WITH
REDUCED BRANCHING COMPRISING THIS HETEROLOGOUS
SPBRANCHED1A GENE

5 The present invention falls within the section of human necessities, in the field of agriculture; new plants or processes for obtaining them by means of biotechnology; flowering angiosperms plants, and specifically tomato plants. In particular, the present invention relates to the *SpBRANCHED1a* gene of *Solanum pennellii*, and tomato plants comprising it heterologously, wherein
10 these tomato plants have fewer and/or shorter axillary shoots than a control plant.

BACKGROUND ART

15 A central question in Biology is how the molecular evolution of genomes has led to the great variety of morphologies found in living organisms. Duplication and divergence of developmental genes is thought to have played a key role in the emergence of new traits. Genome-wide phylogenetic studies are helping us elucidate the evolutionary history of duplications in some of these gene families.
20 However, the molecular mechanisms underlying functional divergence following duplication, and the relationship between gene evolution and emergence of morphological novelties are still not well understood.

In angiosperms, branching patterns greatly determine overall plant architecture
25 and affect key aspects of plant life such as nutrient allocation, height, light harvesting efficiency and visibility for pollinators. Branches are generated from meristems formed in the axils of leaves (axillary meristems, AMs) after germination. AMs develop into *axillary buds*, structures comprising short internodes and nodes bearing leaf primordia and further AMs. Buds may remain
30 dormant for long periods of time or undergo internode elongation to generate lateral shoots. Recent studies suggest that axillary bud development is controlled by a conserved genetic network evolved before the radiation of

flowering plants. AM initiation is controlled by Lateral suppressor/*MONOCULM1* genes, Blind/*RAX1* genes, and *LAX PANICLE* genes. Auxin and strigolactone, hormones synthesised in the shoot apex and the root respectively, act as long distance signals to suppress branching. Synthesis and response to strigolactone is controlled by the More axillary branches/Ramosus pathway (MAX/RMS). Genes responding to these signals and acting inside the bud to prevent outgrowth are also conserved: maize *teosinte branched1* (*tb1*) and its orthologs promote bud arrest in monocots (Doebley *et al.* 1997. *Nature*, 386: 485-488; Hubbard *et al.* 2002. *Genetics*, 162: 1927-1935; Hu *et al.* 2003. *J. Plant Physiol. Mol. Biol.*, 29: 507-514; Kebrom *et al.* 2006. *Plant Physiol*, 140: 1109-1117; Takeda *et al.* 2003. *Plant J.*, 33: 513-520). In dicots, duplications of an ancestral *tb1*-like gene generated the *CYC/TB1* clade comprising three types of genes (Howarth and Donoghue 2006. *J. Plant Physiol. Mol. Biol.*, 29: 507-514). One of them includes *BRANCHED1* (*BRC1*) which has retained most of the branch-suppressing activity in *Arabidopsis* (Aguilar-Martinez *et al.* 2007. *Plant Cell*, 19: 458-472). *tb1/BRC1* genes encode transcription factors of the TCP family (Cubas *et al.* 1999. *Plant J.*, 18: 215-222). TCP transcription factors are involved in the control of cell growth and proliferation in meristems and lateral organs, and they have played a key role in the evolution of innovations such as flower zygomorphy.

Despite the general conservation of genes and pathways, a great diversity of branching patterns is found in angiosperms. Moreover, timing of AM initiation, and branch outgrowth in response to environmental and endogenous signals are very divergent among clades. Molecular evolution of genes controlling branch outgrowth must have played an important role in the generation of this diversity. Putative candidate genes to be targets for selection during the evolution of new branching patterns are *tb1/BRC1-like* genes. Indeed, in maize, artificial selection of *tb1* over-expressing alleles was responsible for the strong apical dominance of the domesticated maize plants (Doebley, Stec and Hubbard 1997, Wang *et al.* 1999. *Nature*, 386: 485-488).

BRC1-like genes from *Solanum lycopersicum* (Solanaceae, Asteridae), a dicot species distantly related to *Arabidopsis* (Brassicaceae, Rosidae) and with diverging branching patterns. In contrast to *Arabidopsis*, which has an indeterminate (monopodial) growth habit, tomato, has a determinate (sympodial) growth habit: while in *Arabidopsis* the shoot apical meristem (SAM) grows indefinitely, in tomato, after the production of 8-12 leaves, the SAM is terminated in a cymose inflorescence. Upon flowering, the lateral meristem in the uppermost leaf axil (sympodial bud) is immediately released to give a shoot. This lateral shoot generates several leaves before terminating with another inflorescence, and so on. Additional lateral shoots may grow out from buds of the primary shoot.

In the international patent application WO 2010/081917 A1 was described the sequence of *SIBRC1* from *Solanum lycopersicum* and inventors elucidated, by means of RNAi experiments, that the *SIBRC1b* gene plays a more important role in the control of axillary bud development and branch outgrowth than *SIBRC1a*. Furthermore, this patent application related to the promoter of said gene. Despite the results shown in that document, it remains in the state of the art the problem to produce plants that have effectively a substantial reduction in the number of branches in order obtain plants with better fruits and also to improve the mechanical harvesting of these fruits.

SUMMARY OF THE INVENTION

The invention relates to a new sequence of *BRANCHED1a* gene, the *SpBRANCHED1a* gene of *Solanum pennellii*, and to an expression product of said gene, to a vector, host cell, cell culture, recombinant expression system, or tomato plants comprising said *SpBRANCHED1a* gene. The present invention also refers to the heterologous use of the gene in tomato plants, to produce a plant having fewer and/or shorter axillary shoots than a control plant, and to a method for producing tomato plants having fewer and/or shorter axillary shoots

than a control plant.

In the present invention the term *SpBRANCHED1a* as synonymous of *SpBRC1a* can be used. Along the present invention, to refer to this gene can be used the term "isolated polynucleotide" or "polynucleotide" encoding a polypeptide comprising the amino acidic sequence of the protein *SpBRANCHED1a* (*SpBRC1a*) from *Solanum pennellii*, taking into account that, by alternative splicing, two proteins are translated from a single genomic sequence, a short and a long protein, differing in their C-terminal domain.

As it is known in the state of the art, in the case of tomato *BRC1*-like genes, *SIBRC1b* seems to have retained the ancestral *BRC1*-like gene function in the suppression of shoot branching while inventors have not been able to establish any relevant role in this process for *SIBRC1a*. Consistently, an asymmetrical distribution of selective constraints has been detected in the evolution of *Solanum BRC1a* and *BRC1b* lineages: the coding sequence of *BRC1b* genes have evolved under a strong purifying selection while the *BRC1a* lineage had experienced a decrease of evolutionary constraints. Therefore, despite this evidence, the present invention shows that *SpBRC1a* plays an unexpected role in suppressing shoot branching when is expressed heterologously in tomato, in opposition with the non-contributing role to this branchless phenotype of the native gen *SIBRC1a*.

The distinguishing features of the present invention in respect to the relevant background art is the new sequence *SpBRC1a* from *Solanum pennellii* and its use to produce plants which express *SpBRC1a* heterologously in tomato plants, as for example, in *Solanum lycopersicum* plants. The technical effect resulting from the distinguishing features is a stronger apical dominance, an improved architecture and more reduced number of lateral branches (branchless phenotype) than control plants, M82 tomato plants. This technical effect implies an improvement of the described phenotype with respect to control plants, described in the examples of the present invention (e.g. IL3-5 plants). This

improvement is due to the functional substitution of the native gene *SIBRC1a* from *Solanum lycopersicum*. Therefore, the expression of *SpBRC1a* in *Solanum lycopersicum* implies an improvement of branchless phenotype in tomato plants.

5

Then, the objective technical problem solved by the present invention is the production of tomato plants with the maximal reduction in number and length of axillary shoots as possible because lateral shoot branching is an undesired trait in the domesticated tomato as it diverts assimilates away from developing fruits.

10

In conclusion, the present invention provides new plants with branchless phenotype modifying or adapting the closest prior art to provide the described technical effect. Therefore, the present invention provides tools to the state of the art to improve the branchless phenotype in tomato plants.

15

It must be stressed that the present invention emphasizes an unexpected or surprising effect: Since is not *SIBRC1a* but *SIBRC1b* the sole responsible in controlling the shoot branching in *Solanum lycopersicum* plants, the native *SIBRC1a* gene does not play a major role in the suppression of axillary bud outgrowth in these tomato plants. In the state of the art is not suggested that any homologous gene of *SIBRC1a* in any tomato plant would have a different activity/function than it is demonstrate with *SIBRC1a*, despite the high identity between proteins *SpBRC1a* and *SIBRC1a*, being 92-93% identical (table 1), the role of *SpBRC1* is very different when expressed in *Solanum Lycopersicum* than the role of the native protein *SIBRC1a*.

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The plants of the present invention also have the advantage to facilitate the pruning and more important than this, because the plants of the invention don't need the pruning of lateral shoots, fewer wounds can be practice and then, potentially, a less number of pathogens will infect the plant.

30

The present invention is susceptible of industrial application, i.e. for avoid the

deviation of assimilates away from developing fruits and also to improve the mechanical harvesting of tomatoes.

Thus, a first aspect of the present invention refers to an isolated polynucleotide,
5 encoding a polypeptide comprising:

- a. an amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or which is at least 90% identical to SEQ ID NO: 2, or
- b. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1
10 or SEQ ID NO: 2.

In the present invention, the terms “polypeptide” and “amino acidic sequence” are synonymous.

15 The amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or which is at least 90% identical to SEQ ID NO: 2 are isoforms of these sequences or sequences with a maximum of 10% of changes in amino acids between them sequences having substantially the same activity/role in the branchless phenotype than SEQ ID NO: 1 or SEQ ID
20 NO: 2. Therefore, the first aspect of the present invention also refers to an isolated polynucleotide, encoding a polypeptide comprising:

- a. the amino acidic sequence of any protein SpBRC1a from *Solanum pennellii*, wherein the protein SpBRC1a has substantially the same role in the branchless phenotype than SEQ ID NO: 1 and/or SEQ ID NO: 2 in tomato
25 plants, or
- b. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1 or SEQ ID NO: 2.

SEQ ID NO: 1 or SEQ ID NO: 2 are the two resulting sequences of alternative
30 splicing of the gene *SpBRC1a*, coded in the genomic sequence (example 1). Since the putative splicing sites (gt/ag) of intron I were found in all the *Solanum BRC1a* genes studied but not in *BRC1b* genes, inventors demonstrate that this

pattern of alternative splicing of *BRC1a* gene was maintained in respect to *SpBRC1a* gene identifying said intron I in the genomic sequence of *SpBRC1a* gene. Then, these two sequences SEQ ID NO: 1 and SEQ ID NO: 2 are involved in the branchless phenotype. The two sequences have different C-terminal (C-t) domains due to a frameshift caused by the alternative splicing of intron I, like in *SIBRC1a* gene.

A preferred embodiment refers to the isolated polynucleotide wherein said polynucleotide is SEQ ID NO: 3 encodes SEQ ID NO: 1, or SEQ ID NO: 4 encodes SEQ ID NO: 2. The sequence SEQ ID NO: 3 is the Coding Sequence (CDS) of the short variant of *SpBRC1a* protein; SEQ ID NO: 1. The sequence SEQ ID NO: 4 is the CDS of the long variant of *SpBRC1a* protein; SEQ ID NO: 2.

In another preferred embodiment of the present invention, the isolated polynucleotide is the genomic sequence SEQ ID NO: 5. This sequence has the introns described for *SIBRC1a*.

The term “% identical” as is understood in this invention refers to the % of identity between two amino acid sequences. The % of identity is a count of the number of positions over the length of the alignment of two sequences where all of the amino acids at that position are identical.

Preferably, the amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or to SEQ ID NO: 2 can be at least 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% identical to SEQ ID NO: 1 or to SEQ ID NO: 2. Furthermore, the amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or which is at least 90% identical to SEQ ID NO: 2 can be an amino acidic sequence with at least less than 90% identical to SEQ ID NO: 1 or SEQ ID NO: 2 if said amino acidic sequence has substantially the same role in the branchless phenotype of SEQ

ID NO: 1 and/or SEQ ID NO: 2 in tomato plants, and also is a sequence from *Solanum pennellii*.

In order to yield information about of the identity of different sequences of BRC1a proteins in tomato plants, table 1 and table 2 is presented.

Table 1. Percentage of identity of several homologous amino acidic sequences respect to SEQ ID NO: 1 or SEQ ID NO: 2 from different tomato plants.

Species 1	Size (amino acids)	Species 2	Size (amino acids)	Identity (%)
<i>S. pennellii</i> Short (SEQ ID NO: 1)	313	<i>S. lycopersicum</i> Short (SEQ ID NO: 8)	325	92
		<i>S. tuberosum</i> Short (SEQ ID NO: 9)	300	85
<i>S. pennellii</i> Long (SEQ ID NO: 2)	349	<i>S. lycopersicum</i> Long (SEQ ID NO: 10)	346	93
		<i>S. tuberosum</i> Long (SEQ ID NO: 11)	336	86

10

The methods to compare the sequences obtaining identity percentages are known in the state of the art, and include, but not limited to, the program GAG, including GAP (Devereux *et al.* 1984. Nucleic Acids Research, 12: 287), Genetics Computer Group University of Wisconsin, Madison, (WI); BLAST, BLASTP, ClustalW2 or Phylemon web server (Tarraga *et al.* 2007. Nucleic Acids Res, 35: W38-42). The percentage of identity has been chosen according to the ClustalW2 Multiple Sequence Alignment from European Bioinformatics Institute.

15

As it is shown in table 1, despite the high identity between proteins SpBRC1a and SpBRC1a, being 92-93% identical, the role of SpBRC1a very different, as demonstrated in the present invention, when expressed in *Solanum*
 5 *Lycopersicum* compared with the role of the native protein SIBRC1a.

Table 2. Percentage of identity between SEQ ID NO: 1, SEQ ID NO: 2 (SpBRC1a) and SpBRC1b from *Solanum pennellii* tomato plants.

Species 1	Size (amino acids)	Species 2	Size (amino acids)	Identity (%)
<i>S. pennellii</i> Short (SEQ ID NO: 1)	313	<i>S. pennellii</i> Long (SEQ ID NO: 2)	349	92
<i>S. pennellii</i> Short (SEQ ID NO: 1)	313	<i>S. pennellii</i> SpBRC1b (SEQ ID NO: 10)	362	37
<i>S. pennellii</i> Long (SEQ ID NO: 2)	349	<i>S. pennellii</i> SpBRC1b (SEQ ID NO: 10)	362	42

10

A preferred embodiment of the present invention refers to the polynucleotide operatively linked to any regulatory sequence that controls the expression of said polynucleotide. In further preferred embodiment of the present invention, said regulatory sequence is SEQ ID NO: 6, SEQ ID NO: 7 or CaMV35S.

15

The term "regulatory sequence that controls the expression of a polynucleotide" refers to a nucleotidic sequence having an effect on the functionality of the polynucleotide gene referred to the beginning of the transcript from a DNA sequence or at the start of translation of a sequence of RNA or other sequences
 20 not described. By way of example, between regulatory sequences of the gene

expression referred to in the present invention are the promoters and other less common as certain introns. The nature of these regulatory sequences differs depending on the host organism; in prokaryotes, these sequences of control usually include a promoter, a site of union ribosomal, and signs of completion; in eukaryotes, usually, these sequences of control include promoters, signs of termination, intensifiers and sometimes silencers.

As it is used here, the term "promoter" refers to a region of the DNA upstream from the start point of the transcription of a gene, and particularly, that is able to start the transcription in a plant cell. Examples of promoters include but are not limited to, promoters obtained from plants, virus of plants or bacteria that can express genes in cells of plants, as *Agrobacterium* or *Rhizobium*. Promoters can be classified, for instance, as inducible and constitutive promoters. The sequence SEQ ID NO: 6 is the inducible promoter of the gene *SIBRC1a* from *Solanum lycopersicum*. The sequence SEQ ID NO: 7 is the inducible promoter of the gene *SIBRC1a* from *Solanum tuberosum*.

The term "operatively linked" refers to a juxtaposition in which the components as well described have a relationship that enables them to work in the intentional way. A polynucleotide operatively linked to any regulatory sequence that controls the expression of said polynucleotide is linked in such a way that the expression of the coding sequence of the polynucleotide is achieved under conditions compatibles with the regulatory sequence.

To refer to any of the polynucleotide described in the above aspect and its preferred embodiments of the present invention, the term "polynucleotide of the invention" or "polynucleotide of the present invention" can be used.

Another aspect of the present invention refers to an expression product of the polynucleotide of the invention. To refer to any of the expression product of the polynucleotide of the present invention, the term "expression product of the invention" or "expression product of the present invention" can be used.

The term "product of the expression of the sequence" as it is used in the present invention refers to any product derived from the expression of the polynucleotide of the invention. Thus, as product derived from the expression of the polynucleotide of the invention is understood, for example, the RNA that is obtained from the transcription of said polynucleotide, the processed RNA, the resulting protein in the translation of RNA, or subsequent modifications to the polynucleotide inside the cell, provided that the resulting sequence of these modified polynucleotides having its origin in the original sequence or the functional role described in the present invention.

A further aspect of the present invention refers to an expression vector comprising the polynucleotide of the invention. To refer to any vector comprising the polynucleotide of the present invention, the term "vector of the invention" or "vector of the present invention" can be used.

The term "vector" refers to a DNA fragment that has the ability to replicate in a given host and, as the term indicates, it can serve as a vehicle to multiply another fragment of DNA that has being linked to the same (insert). The term "insert" refers to a DNA fragment that is linked to the vector; in the case of the present invention, the vector comprises any of the polynucleotides of the invention that can be replicated in the appropriate host. The vectors may be plasmids, cosmids, bacteriophages or viral vectors, without excluding another type of vectors that correspond to this definition of vector.

Another aspect of the present invention refers to a host cell comprising the polynucleotide, the expression product, or the vector of the invention, wherein if the host cell is a plant cell, said plant cell is not from the species *Solanum pennellii*. That is to say, the polynucleotide of the present invention (the mean core corresponding to the gen of *Solanum pennellii*) is present in said host cell in heterologous form. A preferred embodiment refers to the host cell of the

present invention, wherein said cell is a plant cell. More preferably the plant cell is a *Solanum lycopersicum* cell.

5 The term "host cell" as it is used in the present invention relates to any procariotic or eukaryotic cell, essentially, it refers to a eukaryotic plant cell and within this group, more preferably, those cells belonging to the Kingdom Plantae, wherein any of these cells comprises polynucleotide, the expression product, or the vector of the invention. Thus, the term plant cell comprises at least one cell of the parenchyma, meristematic cell or of any kind of plant cell,
10 differentiated or undifferentiated. Preferably the plant cell is a tomato plant cell and more preferably is a *Solanum lycopersicum* cell.

To refer to any host cell comprising the polynucleotide of this invention, the term "host cell of the invention" or "host cell of the present invention" can be used.

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The present invention further relates to a cell culture comprising the host cell of the invention. To refer to any cell culture comprising the host cell of the present invention, the term "cell culture of the invention" or "cell culture of the present invention" can be used.

20 The term "cell culture" refers to a cultivation of cells isolated from the same or different type of tissue, or a collection of these cells organized in parts of a plant or in tissues (tissue culture). Types of this kind of cultures are, e.g. but without any limitation, a culture of protoplasts, calli (groups of plant cells undifferentiated able to regenerate a complete plant with the appropriate
25 organogenic program) or a culture of plant cells that are isolated from plants or parts of plants such as embryos, meristematic cells, pollen, leaves or anthers.

Another aspect of the present invention refers to a recombinant expression system comprising the polynucleotide, the expression product, the vector, or the
30 host cell of the invention.

The recombinant expression system can be, but without limitation, e.g. a recombinant host cell or a recombinant bacteriophage or their combination with any helper virus. Recombinant manufacturing involves the expression of a DNA construct encoding for the desired protein in a recombinant host cell. The host cell can be either prokaryotic or eukaryotic. Recombinant manufacturing, however, does have its difficulties. Expression constructs must be optimized for a particular protein and for a particular host cell. Expressing a recombinant protein in a host cell exposes the recombinant protein to a new set of host cell enzymes, such as proteases, which can modify or even degrade the recombinant protein. Modification and degradation of the recombinant protein of interest is undesirable.

A further aspect of the present invention refers to a use of the polynucleotide, the expression product, the vector, the host cell, the cell culture, or the recombinant expression system of the invention, to produce a tomato plant, but not a plant of the species *Solanum pennellii*, having fewer and/or shorter axillary shoots than a control plant.

The term "axillary shoots" can be used as synonymous of "lateral branches", "lateral shoots", "secondary shoots", "axillary shoot branching", "axillary bud outgrowth", and refers to a shoot growing from the axillary bud located between the leaf and the stem, that is to say, growing in a node. In a control plant, the stem displays branching due the outgrowth of axillary buds in the individual node positions.

In the tomato plants of the present invention, the length of the basal lateral branches (nodes 1-6) presents significant differences with regard to the length of the lateral branches in the control.

In the present invention, the term "fewer and/or shorter" relates to significant differences between the presence/absence and/or the length of axillary shoots, respectively, in the equivalent node of the plants of the invention and control

plants. The skilled person can arrive to these significant differences using any statistical and any method known in the background art. Then, in the present invention plants having fewer and/or shorter axillary shoots than a control plant have branchless phenotype.

5

For instance, as shown in the examples of the present invention, IL3-5 plants had significantly fewer and/or shorter axillary shoots than M82 plants in cotyledons, and nodes 1, 4, 5 and 6.

- 10 The control plant is any tomato plant, wild type carrying empty vectors, but in any case the control plant does not contain the polynucleotide of the present invention. Preferably the control plant is a tomato plant belonging to the species *Solanum lycopersicum*, preferably the control plant is a M82 cultivar of tomato. The control plant has the same taxonomic category than the tomato plant
- 15 comprising the polynucleotide of the invention in order to compare the number or/and length of lateral branches.

- Another aspect of the present invention refers to a tomato plant, but not a *Solanum pennellii* plant, comprising the polynucleotide, the expression product,
- 20 the vector, the host cell, the cell culture, or the recombinant expression system of the invention, wherein said plant has fewer and/or shorter axillary shoots than a control plant. Thus, the polynucleotide of the present invention (the mean core corresponding to the gen of *Solanum pennellii*) is present in plant in heterologous form. Preferably the tomato plant belongs to the species *Solanum*
- 25 *lycopersicum*.

- The tomato plant of the present invention can be selected from potentially tomato plant species, for instance the tomato plant can be selected from tomato plants with red, yellow, orange or yellow-green fruit, and without limitation to
- 30 these species: *Solanum lycopersicum*, *Solanum peruvianum*, *Solanum pimpinellifolium*, *Solanum lycopersicoides*, *Solanum sitiens*, *Solanum*

juglandifolium, *Solanum ochranthum*, *Solanum cheesmaniae* or *Solanum galapagense*.

Preferably, the tomato plant belongs to the species *Solanum lycopersicum*.

- 5 More preferably the variety of the tomato plant can be selected from the list of stock names in the SOL Genomics network (SGN) Breeders Toolbox (<http://solgenomics.net/breeders/index.pl>). Preferably the variety of the tomato plant from the species *Solanum lycopersicum* can be selected from the list, but without limitation, Anna russian, Applause, Aussie, Baladre, Bella rosa, Black
- 10 cherry, Black russian, Blondkopfchen, Brandywine, Carbon, Ceylan, Cherokee purple, Cherry yellow pear, Black Plum, Comanche, Copy, Costoluto genovese, Ditmarcher, Eros, Gallician, Glacier, Gartenperle, Green sausage, Grushovka, Harzfeuer, Hugh, Japanesse black, Jersey devil, Kosovo, Krim black, Kumato, Liguria, Limachino, Lime green salad, Manitoba, Marvel stripe, Moneymaker,
- 15 Muchamiel, Opalka, RAF, Black Pear, Hawaian pineapple, Rio grande, San marzano, Siberian, Sprite, Sugary, Sun sugar, Tigerella, Valencian, White Queen or Window box Roma.

A further aspect of the present invention refers to a tomato plant comprising, heterologously, a fragment of the chromosome III of *Solanum pennellii* that comprises a polynucleotide encoding a polypeptide comprising:

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- a. an amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or which is at least 90% identical to SEQ ID NO: 2, or
- b. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1 or SEQ
- 25 ID NO: 2.

,wherein said plant has fewer and/or shorter axillary shoots than a control plant

A preferred embodiment of the invention refers to said tomato plant comprising, heterologously, a fragment of the chromosome III of *Solanum pennellii* that comprises a polynucleotide encoding a polypeptide, wherein said fragment of

30 the chromosome III corresponds to the fragment comprised between the DNA marker TG284A and the DNA marker TG244 (figure 2). Preferably the fragment

of the chromosome III corresponds to the fragment comprised between the DNA marker T0794 and the DNA marker TG244; or more preferably to the fragment comprised between the DNA marker TG377 (or the marker TG126 or CD4A) and the DNA marker TG244. The figure 2 shows all the proposed DNA
5 markers within chromosome III of *Solanum pennellii*. The cited markers can be analyzed in detail in the Sol Genomics Network (SGN) database.

A preferred embodiment of the invention refers to the plant comprising, heterologously, a fragment of the chromosome III of *Solanum pennellii*, wherein
10 comprises a polynucleotide encoding a polypeptide comprising SEQ ID NO: 3 encoding SEQ ID NO: 1, or SEQ ID NO: 4 encoding SEQ ID NO: 2. Preferably said polynucleotide is the genomic sequence SEQ ID NO: 5.

A preferred embodiment of this invention refers to any plant described above
15 wherein the expression of the polynucleotide of the invention is greater than the expression of the homologous native gene *BRC1a*.

The plant contains the polynucleotide of the invention in homozygosis, heterozygosis or hemicigosis. A further embodiment of the present invention
20 refers to any plant described above, wherein the polynucleotide is integrated in their genome, preferably integrated in homozygosis

Another aspect refers to a germplasm of the plant of the present invention. Preferably the germplasm is a seed or pollen.
25

The term "plant of the present invention" includes each of the parties on said plant, which can be conserved or cultivated in isolation or in combination, as well as the germplasm. The germplasm is defined by the biological material that contains the intraspecific genetic variability or by the genetic materials that can
30 perpetuate a species or a population of said plant. Thus germplasm is the seed, tissue culture for any part of the plant or plants established in *ex situ* collections, without excluding any other material in the scope of this definition.

The pollen has high level of interest since the transmission of the genetic and phenotypic characters can be carried out by the pollination of any plant variety compatible with the pollen that is referenced. In this way can be produce a plant
5 which includes the polynucleotide of the invention, after the respective cross and selection, it can be obtained a plant in which the polynucleotide integrates a suitable number of copies in stable condition in order to obtain the same desirable branchless phenotype in the subsequent generations.

10 To refer to any of the tomato plants of the present invention, the term “plant of the invention” or “plant of the present invention” can be used.

The plant of the invention does not belongs to the species *Solanum pennellii*, then being the polynucleotide of the present invention expressed in different
15 plant species that *Solanum pennellii*, therefore, the polynucleotide is expressed heterologously.

A further aspect of the present invention refers to a method for producing tomato plants having fewer and/or shorter axillary shoots than a control plant,
20 comprising:

- a. Transforming at least a tomato plant cell with a heterologous polynucleotide comprising:
 - i. an amino acidic sequence from *Solanum pennellii* which is at least 90% identical to SEQ ID NO: 1 or to SEQ ID NO: 2, or
25 ii. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1 or SEQ ID NO: 2, and
- b. growing the plant cell obtained in the step (a) in a suitable medium to produce at least a plant which expresses the heterologous polynucleotide.

30 Another embodiment of the present invention refers to the method for producing tomato plants having fewer and/or shorter axillary shoots than a control plant, wherein said tomato plant belongs to the species *Solanum lycopersicum*.

The plant of the invention can be achieved by genetic transformation mediated by biolistic, *Agrobacterium tumefaciens* or any other technique known by the skilled in the art (e.g. transformation of protoplasts), that will allow the
5 integration of the polynucleotide of the invention in any of the DNA of the plant; genomic, chloroplastic or the polynucleotide of the invention by crossing and selection.

The suitable medium growing the plant cell obtained in the step (a) is a known
10 medium by the skilled in the art.

The production of a plant as is indicate in step (b) can be performed by means techniques known by the skilled in the art, for instance:

- The cultivation of embryos: Isolation of zygotic embryos promoting their growth as plant in an artificial environment
- 15 - Somatic embryogenesis: Production of embryos from somatic tissues, such as microspores or leaves
- Organogenesis: Production of organs such as stems or roots from various tissues of the plant.

20 The plant of the present invention can be obtained according with other microbiologic processes as for example:

- Obtaining cybrids: It is produced a cell with its cytoplasm and the cytoplasm of the other cell and this cell can be grown in a suitable medium to produce a plant
25 which expresses the heterologous polynucleotide.

- Fusion of somatic cells (preferably protoplast fusion): at the cytoplasmic level hybrid plants can be the result of: (a) The sum of the cytoplasm of both parental; (b) The cytoplasm of a single parental; (c) a cytoplasm hybrid result of
30 recombination of the genomes extranuclears of both cells. The fusion of somatic cells can be applied; to overcome the incompatibility in interspecific crosses, or

a better utilization of the interspecies variation and extraspecific in interspecific compatible crosses.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Methods and materials similar or equivalent to those described herein can be used in the practice of the present invention. Throughout the description and claims the word "comprise" and its variations are not intended to exclude other technical features, additives, components, or steps. Additional objects, advantages and features of the invention will become apparent to those skilled in the art upon examination of the description or may be learned by practice of the invention. The following examples, figures and sequence listing are provided by way of illustration and are not intended to be limiting of the present invention.

15

BRIEF DESCRIPTION OF THE FIGURES

In order to complement the description that is being made and in order to help better understanding of the characteristics of the invention and in agreement with some preferred examples, figures have been included where, with illustrative and non-limiting character, the following is represented:

20

FIG. 1. Shows the *BRC1*-like genes and proteins in tomato.

(a) Maximum Likelihood (ML) phylogenetic tree with 1000 bootstrap pseudoreplicates of class II TB1/CYC genes from tomato (Sl, bold) and representative class II TCP members from *Antirrhinum majus* (Am), *Arabidopsis thaliana* (At), *Solanum tuberosum* (St) and *Populus trichocarpa* (Pt). Branches with support of 500 or more are indicated. AtTCP4/CIN are the outgroups. (b) Gene structure of *SIBRC1a* and *SIBRC1b*. Coding sequences are shaded in black, introns in white, 5' UTR and 3'UTR in light grey. Below each gene, predicted proteins encoded by the isolated cDNAs are represented. Blue boxes

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indicate TCP and R domains. Red boxes indicate the alternative coding frame in the short SIBRC1a protein. Dark grey box indicates the alternative intron translated in the long protein.

5 **FIG. 2. Shows the map of chromosome III of *Solanum lycopersicum* (a) and *Solanum pennellii* (b) and the relative position of *BRC1* gene.**

In both maps are indicated several DNA markers and their relative position between them and between BCR1a gene and IL3-5 introgressed fragment.

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TG479, T1286, CT171, T1511, T0794, TG244 are DNA markers from chromosome III of *Solanum lycopersicum* and TG40, CAB3, TG13S, CT31, TG114, TG585, TGS17, TG130, TG13B, CT22, TG66, TG39, TG247, TG288, TG50C, TG102, CT90A, TM8, TGS99, TG246, TG457, TG42, TG134, TG284A, TG377, CD4A, TG126, TG152, CD71, TG406B, TG214, CD69, CT250, TG244 and TG94. The fragment of the Chromosome III from *Solanum Pennellii* introgressed in *Solanum lycopersicum* in IL3-5 introgression line is shown with the bar IL 3-5. (a) and (b) shows some common DNA markers in both chromosomes.

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FIG. 3. Shows the Secondary structure prediction of the long (top) and short (bottom) C-t motifs of SIBRC1a according to Jpred.

E, extended secondary structure, H, α -helix. Jnet, solvent accessibility predictions.

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B, buried, -, exposed. Jnet Rel, reliability of prediction accuracy, ranges from 0 to 9.

FIG. 4. Shows the analysis of *SIBRC1a* function.

(a) Shoot branching phenotype of *Sibrca-v171i* point mutants compared to their wild type siblings. Lateral branch length of mutant tomato plants and their wild type siblings at anthesis of the first flower. Standard errors (\pm s.e.m.) are represented.

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$N_{brc1a-v171i}=7$, $N_{wt}=25$. (b) Shoot branching phenotype of IL3-5 and M82 plants

measured as in (a) $N_{IL3-5}=12$, $N_{wt\ M82}=15$. Asterisks indicate significant differences ($p<0.05$). (c) Young M82 wild type (left) and IL3-5 (right) plants grown in the same conditions. Arrows indicate branches grown out in wild type plants and arrested buds in IL3-5. (d). Relative *SIBRC1a* mRNA levels in axillary buds of M82 and IL3-5 plants. The asterisk indicates a significant difference ($p<0.05$).

FIG. 5. Shows the phenotype of adult M82 and IL3-5 plants after flowering.

10 EXAMPLES

The following examples provide a description, of illustrative and non-limiting character, of some of the assays and operating conditions claimed in the list given below that refers to the cloning and use *SpBRC1a* as a suppressor of shoot branching in tomato, as well as plants of *Solanum lycopersicum* expressing the *SpBRC1a* gen.

In the present examples the term *SpBRC1a* is equivalent to an isolated polynucleotide encoding a polypeptide comprising the amino acidic sequence of SEQ ID NO: 1 (short splicing alternative protein) or SEQ ID NO: 2 (long splicing alternative protein). *SIBRC1a* and *SIBRC1b* refer to the gen *BRANCHED1a* or *BRANCHED1b* of *Solanum lycopersicum*.

25 EXAMPLE 1. Identification of tomato *BRC1*-like genes

The inventors searched for *CYC/TB1* type genes in tomato genomic and EST databases (Mueller 2009. The Plant Genome, 2: 78-92) and identified six TCP genes of this group. Phylogenetic analyses showed that two were *CYCLOIDEA* (*CYC*)-like genes, two were *BRC2*-like genes and another two were *BRC1*-like genes. These six TCP genes were named *SICYC1*, *SICYC2*, *SIBRC2a*, *SIBRC2b*, *SIBRC1a* and *SIBRC1b* (*Solanum lycopersicum* *CYC*, *BRC2*, *BRC1*) respectively (Figure 1a) and were mapped to tomato chromosomes 2-6 (Figure 2). *CYC*-like

genes have been shown to be expressed in floral meristems and to be mainly involved in the control of flower shape in many species. Inventors focussed on the two *BRC1*-like genes (*SIBRC1a* and *SIBRC1b*) and investigated their roles during tomato lateral shoot development.

5

Inventors first studied the transcriptional activity of these genes. For *SIBRC1a*, inventors isolated two cDNAs generated by alternative splicing: a short one, with a 978 base pairs (bp) long open reading frame (ORF) split by two introns (introns I and II), and a long one (1041 bp), in which intron I was retained and translated (Figure 1b). Splicing of intron I was a relatively rare event as long cDNAs were isolated 9 times more frequently than short cDNAs. These cDNAs encoded two proteins with identical N-terminal, TCP and R domains but different C-terminal (C-t) domains of 36 and 57 amino acids respectively, due to a frameshift caused by the alternative splicing of intron I (Figure 1b). For *SIBRC1b*, inventors isolated a single type of cDNA containing a 1089 bp long ORF split by one intron (Figure 1b).

10
15

Alignment of the short and long cDNAs of *SIBRC1a*, the cDNA of *SIBRC1b* and the *SIBRC1a* and *SIBRC1b* genomic sequences confirmed that intron II of *SIBRC1a* was co-linear with the intron of *SIBRC1b*, suggesting that this intron was also spliced in the common ancestor, before duplication. In contrast, intron I of *SIBRC1a* could have evolved later, after duplication: it had frame conservation and a high sequence similarity to a coding (unspliced) region of *SIBRC1b*. Moreover, the C-t peptide encoded by the long *SIBRC1a* cDNA (which retained intron I) had sequence similarity to the C-t peptide of *SIBRC1b*, supporting that this frame was translated in the common ancestor, which did not splice this region. The putative splicing sites (gt/ag) of intron I were found in all the *Solanum BRC1a* genes studied but not in *BRC1b* genes, suggesting that they evolved within the *BRC1a* clade. Interestingly, the C-t region of the short *BRC1a* protein was predicted to form a strongly amphipathic α -helix (Figure 3) highly conserved in all the *Solanum* species analyzed. On the other hand the C-t peptide of the long *SIBRC1a* protein was predicted to have an extended secondary structure (Figure 3) like that of *SIBRC1b*. These results indicate that, after the separation of Arabidopsis and Solanaceae, a

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duplication of an ancestral *BRC1*-like gene, which had one intron, generated two gene copies, *BRC1a* and *BRC1b*. A new alternative splicing site evolved in the *BRC1a* clade generating a divergent transcript, with a 3'-terminal frameshift, encoding a protein with a novel C-t motif predicted to form a strongly amphipathic α -helix.

Since the putative splicing sites (gt/ag) of intron I were found in all the *Solanum BRC1a* genes studied but not in *BRC1b* genes, inventors followed this pattern of alternative splicing of *BRC1a* gene in the genome sequence of *SpBRC1a* gene and, after was identified said intron I, inventors predicted the short cDNA coding for SEQ ID NO: 1 and the long cDNA coding for SEQ ID NO: 2.

EXAMPLE 2. *SpBRC1a* acts as a suppressor of shoot branching in tomato
Role of *SIBRC1a* and *SIBRC1b*.

The finding of two tomato *BRC1*-like paralogs more closely-related to each other than with *AtBRC1*, suggests that a duplication of *BRC1* took place after the separation of the Brassicaceae and Solanaceae. These two *BRC1*-like genes map to different chromosomes, suggesting that they did not originate by segmental duplication but rather by chromosomal duplication or during the whole-genome duplication (WGD) occurred in Solanaceae 60–70 mya. Duplication, a common event in the evolution of genomes, is usually followed by loss of one of the gene copies by accumulation of deleterious mutations. In Arabidopsis, for instance, a single *BRC1* gene has been found, although this species has undergone several WGD suggesting that all the other *BRC1*-like gene copies have been lost. A large number of models have been proposed to explain fixation, maintenance and evolution of duplicate genes.

These results indicate that in the case of tomato *BRC1*-like genes, *SIBRC1b* seems to have retained the ancestral *BRC1*-like gene function in the suppression of shoot branching while inventors have not been able to establish any relevant role in this process for *SIBRC1a*. Consistently, an asymmetrical

distribution of selective constraints has been detected in the evolution of *Solanum BRC1a* and *BRC1b* lineages: the coding sequence of *BRC1b* genes have evolved under a strong purifying selection while the *BRC1a* lineage had experienced a decrease of evolutionary constraints.

5

2.1. Role of *SIBRC1a* and *SIBRC1b* in tomato and its evolution in *Solanum* genus.

As far as it is known, *SIBRC1b* acts as a suppressor of shoot branching in tomato whereas *SIBRC1a* does not play major role in the control of shoot
10 branching in tomato (WO 2010/081917 A1). In the above-mentioned patent application inventors arrived to this conclusion after carrying out experiments using RNAi for each of the genes *SIBRC1a* and *SIBRC1b*, taking this fact into account in the final claims, because neither the use of *SIBRC1a* nor the use of *SIBRC1a*-RNAi contributed to solve any technical problem. In the present
15 application inventors repeated the experiments of silencing and the same results that it is shown in WO 2010/081917 were obtained; *SIBRC1b* loss-of function leads to increased branch outgrowth in tomato but *SIBRC1a* loss-of function does not cause bud outgrowth. As a result, these examples are excluded in the present application because they do not provide more relevant
20 information.

2.1.1. Role of *SIBRC1a*

The *Solanum lycopersicum* wild-type branching patterns of strong *Slbrc1a-v220i*
25 mutants could indicate that *SIBRC1a* does not play a major role in the suppression of axillary bud outgrowth in tomato, raising the possibility that this gene has become non-functional in this species. Other signs of pseudogenization such as the accumulation of missense mutations have not been found in this tomato gene. Moreover, transcription is tightly regulated
30 spatially and temporally, and alternative splicing takes place, indicating that transcriptional and post-transcriptional regulation have not been lost. No signs of a trend towards pseudogenization have been detected in the evolution of the

Solanum BRC1a lineage, either. These results indicate that the fast evolution rate found in this clade is not due to relaxation (usually associated with loss of function), but rather to positive selection and adaptation. Moreover, inventors have identified several protein sites, targets of positive selection and evolving at high evolution rate, located in domains potentially involved in stability (PEST domain) and transcriptional activity, aspects which could greatly affect protein function. In addition, the alternative splicing evolved within the *BRC1a* clade gives rise to a novel transcript (identified both in tomato and potato) encoding a divergent protein with a C-t amphipathic helix, secondary structure often involved in protein-protein interactions, dimerization and transcriptional activation or repression. Inventors speculate that this novel domain, generated by a 3' terminal (3'-t) frameshift of the *BRC1a* transcripts, could also be contributing to the structural (and perhaps functional) divergence of *BRC1a* proteins. 3'-t frameshift mutations have been proposed to be responsible for the evolution of novel protein domains, instrumental to the diversification of transcription factor families. In summary, the *BRC1a* clade could be diversifying rather than leading to loss of function and pseudogenization.

In the particular case of the tomato *SIBRC1a* gene, its low transcription levels and the apparently irrelevant function could be result of the recent artificial evolution associated to tomato domestication. Under-expressor alleles could have been fixed during the selection of other linked loci controlling selected traits. Alternatively, *SIBRC1a* could be functional but only up-regulated under certain developmental or environmental conditions not yet identified. So far, inventors have not detected up-regulation of *SIBRC1a* in plants with reduced *SIBRC1b* function, for instance, but other genetic or environmental situations might lead to transcriptional activation of *SIBRC1a*.

2.1.2. Role of *SIBRC1b*

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The excess of shoot branching displayed by *SIBRC1b* loss-of-function plants resembles that of *tb1/brc1* mutants in other species (Aguilar-Martinez *et al.*

2007. *Plant Cell*, 19: 458-472; Doebley *et al.* 1997. *Nature*, 386: 485-488; Finlayson 2007. *Plant Cell Physiol.*, 48: 667-677; Minakuchi *et al.* 2010. *Plant Cell Physiol*, 51: 1127-1135, Takeda *et al.* 2003. *Plant J.*, 33: 513-520) and indicates that *SIBRC1b* must have retained the ancestral role in branch growth suppression. This is consistent with the low evolution rate (w) of the *BRC1b* genes in *Solanum*. *SIBRC1b* cis-regulatory regions and trans-acting factors also seem to be conserved: *SIBRC1b* is mainly expressed in arrested axillary buds, especially in basal nodes, like *AtBRC1* and rice *fine culm1*, and it responds to decapitation like *AtBRC1* (Aguilar-Martinez *et al.* 2007, Arite *et al.* 2007. *Plant Journal*, 51: 1019-1029). In *Arabidopsis* and rice it has been proposed that *AtBRC1* and *FC1*, respectively, are downstream of the strigolactone pathway (Aguilar-Martinez *et al.* 2007. *Plant Cell*, 19: 458-472; Minakuchi *et al.* 2010. *Plant Cell Physiol*, 51: 1127-1135). Although in tomato this needs to be tested, loss of function of *SICCD7*, a gene involved in tomato strigolactone synthesis (Vogel *et al.* 2009. *Plant J*, 61, 300-311; Koltai *et al.* 2010. *J Exp Bot*, 61: 1739-1749) causes alterations in vegetative but not in sympodial branching, similarly to loss of *SIBRC1b* function. However, *Slccd7* phenotypes are much stronger than those of *SIBRC1b* RNAi lines suggesting that additional factors may be controlled by this gene. Other related TCP genes such as *SIBRC1a*, *SIBRC2a* and *SIBRC2b*, are expressed in axillary buds at much lower levels than *SIBRC1b* but inventors cannot rule out that, in some conditions, they are relevant for this process.

2.2. Use of *SpBRC1a* as a suppressor of axillary shoot branching in tomato

Different tomato seed stocks obtained from the Tomato Genetics Resource Centre (TGRC) at the University of California Davis were grown. Special mention can be done with respect to the introgression line IL3-5. As it is indicated in the "chromosomal mapping" section, mapping was done in a *Solanum pennellii* introgression population using primers combinations and, after detection of the corresponding PCR fragments, the inventors concluded

that *BRC1a* mapped to IL3-5 (Figure 2), in addition *BRC1b* mapped to IL6-2 and *BRC2a* to IL4-1. Therefore, the introgressed fragment from *Solanum pennellii* contains *SpBRC1a* gene. For these plants IL3-5, neither “branchless” phenotype nor the expression of *SpBRC1a* has been mentioned to date.

5

Interestingly, as shown in figure 4, IL3-5 lines, carrying a genomic region comprising a *SpBRC1a* gene, expressed this gene, *SpBRC1a*, at four times higher levels than *SIBRC1a* is expressed in M82 control plants and they display stronger apical dominance than M82 plants. These results demonstrate that introgression of expressed wild *SpBRC1a* genes into production tomato lines generates plants with improved architecture and reduced lateral shoot branching (figures 4 and 5).

10

EXAMPLE 3. *SpBRC1a* is the responsible for the strong apical dominance of introgression line IL3-5 plants and also of *SpBRC1a* *Solanum lycopersicum* transgenic plants.

15

To confirm that *SpBRC1a*, contained in the introgression line IL3-5, is the gene responsible for the strong apical dominance of these lines, inventors caused silencing of *SpBRC1a* by generating RNAi transgenic lines carrying a sequence that is identical to part of its coding sequence. Inventors compared the branching phenotype of transgenic plants with those of M82 and with those carrying an empty binary vector. *SpBRC1a* was the responsible for the branch suppression RNAi lines having a similar number of branches as M82 plants.

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25

3.1. *SpBRC1a* and the strong apical dominance of introgression line IL3-5 plants.

3.1.1. CaMV35S:RNAi Constructs

30

A *BRC1a*-specific PCR product (225 bp) was cloned into the vector pHANNIBAL (CSIRO) using restriction sites BamHI/ClaI and XhoI/KpnI for the first and second cloning, respectively. Primers used were A and B:

A: 5'GGGGCTCGAGGGATCCTGGCTACTCACAAAGTCAAATCAGCG 3'
(SEQ ID NO: 17).

B: 5'GGGGGGTACCATCGATAACAACCTGGATGAATTATTGCCCTACG 3'
(SEQ ID NO: 18).

The pHANNIBAL cassettes were digested with NotI and subcloned in the NotI site of Bluescript SK+. The cassette was then digested with SacI/SmaI and cloned in the MCS of the binary vector pBIN19. For the controls inventors will use an empty pBIN19 vector.

3.1.2. Generation of IL3-5 Tomato Transgenic Plants comprising the RNAi Constructs.

Binary vectors were transformed into *Agrobacterium tumefaciens* strain LBA4404. To generate stable transgenic tomato plants inventors will use the *Solanum pennellii* introgression line IL3-5 in the background of *S. lycopersicum* cv. M-82 (LA4060). Inventors transformed cotyledons according to (Ellul *et al.* 2003. Theor Appl Genet, 106: 231-238). Shoots regenerated from calli grown in kanamycin were transferred to non-selective root inducing media and were confirmed for transgene integration by PCR using primers Agri51 and PDK_Hannibal_51.

Agri51: 5'CAACCACGTCTTCAAAGCAA 3' (SEQ ID NO: 19).

PDK_Hannibal_51: 5'ATCTTCTTCGTCTTACACATCACTTG 3' (SEQ ID NO: 20).

Levels of transgene expression were quantified by Real-Time Q-PCR in RNA from leaves. Ploidy level of transformants was analyzed in leaf tissue, using wild type tissue as diploid reference. Nuclei were isolated according to (Galbraith *et al.* 1983. Science, 220: 1049-1051). Cytometry analyses were performed using a flow cytometer Beckman Coulter EPICS XL-MCL. Branch lengths were measured in mm immediately after anthesis of the first flower

3.2. *SpBRC1a Solanum lycopersicum* transgenic plants show branchless phenotype.

5 In addition, in order to confirm that *SpBRC1a* contributes to promote a stronger apical dominance than that of control plants, inventors generated transgenic lines carrying the genomic sequence of *SpBRC1a*; SEQ ID NO: 5. Inventors compared the branching phenotype of transgenic plants with those of M82 control plants and with those carrying an empty binary vector. Inventors
10 observed that transgenic plants have fewer number of branches than M82 plants.

SpBRC1a was cloned in pBIN19 vector under the control of the *SpBRC1a* promoters. Binary vectors were transformed into *Agrobacterium tumefaciens*
15 strain LBA4404 and transgenic plants of *S. lycopersicum* cv. M-82 were generated after regeneration.

EXPERIMENTAL PROCEDURES

20

Plant material

Tomato seed stocks were obtained from the Tomato Genetics Resource Centre (TGRC) at the University of California Davis or kindly provided by R. Fernández
25 Muñoz (Estación Experimental La Mayora, Murcia, Spain). Accession numbers: Moneymaker (MM), LA2706; *Solanum pennellii*, LA0716; *Solanum pimpinellifolium*, PE-2; *Solanum habrochaites*, LA1777; *Solanum galapagense*, LA0317; *Solanum arcanum*, PE-30; *Solanum lycopersicum*, LA2706; *Solanum habrochaites glabratum*, PI 126449; *Solanum huaylasense*, LA2561; *Solanum chilense* (1969), LA1969; *Solanum peruvianum*, LA0098; *Solanum chmielewskii*, LA1028; *Solanum corneliomulleri*, LA0366; *Solanum pennellii*
30 introgression lines in the background of *S. lycopersicum* cv. M-82, IL3-5,

LA4047; LA4060. TILLING *Slbrc1a* point mutant alleles were obtained from tomato M82 EMS-mutagenized population and a Red Setter EMS-mutagenized population. Seeds were germinated in soil and transplanted to 1.5L pots and grown in chambers at 21°C, long photoperiod (16 h light 8 hours dark), PAR 90 mmol m² s⁻¹.

Identification and isolation of tomato *BRC1*-like genes

Inventors carried out BLAST searches at the TIGR Solanaceae Genomics Resource site (http://jcv.org/potato/sol_ma_blast.shtml), TIGR Plant Transcript Assemblies Database (<http://plantta.jcv.org/>) and SOL Genomics Network Tomato Gene Index website (<http://solgenomics.net/>) using the TCP domain of *AtBRC1*. To obtain full length cDNAs, RNA from dissected axillary buds was extracted with the guanidine-HCl method. cDNA was synthesized and full length cDNAs were isolated using primers indicated in Table S2.

Chromosomal mapping of the *CYC/TB1*-like genes

Genomic fragments of *BRC1a*, *BRC1b* and *BRC2a* were amplified from *L. pennellii* and sequenced to find PCR based markers. Mapping was done in an *L. pennellii* introgression population (Eshed and Zamir 1995. Genetics, 141: 1147-1162) using primer combinations TCP1-F2 (5' CCTCATAAAAGGGAATCAAGGGA 3'; SEQ ID NO: 13)/Le3 (5' ATTGAGAATGACTTGAAAGATAAAGATGAG 3'; SEQ ID NO: 14) and the restriction enzyme Csp6I to detect a CAPS for *BRC1a* and TCP3-F2 (5' CCTCATAAAAGGGAATCAAGGGA 3'; SEQ ID NO: 15) /TCP3-R2 (5' TATTGAAAAATCGCGCACGTA 3'; SEQ ID NO: 16) and BspHI for *BRC2a*. A PCR-fragment length polymorphism for *BRC1b* was detected using primers TCP2-F2/LeTCP2 cDNA-R. *BRC1a* mapped to IL3-5, *BRC1b* to IL6-2 and *BRC2a* to IL4-1.

Phenotypic analysis

Branch lengths were measured in mm immediately after anthesis of the first flower.

5 Real-Time PCR

Plant tissue was harvested and RNA was isolated with the guanidine-HCl method. Each biological replicate came from individually dissected axillary buds (Ax buds) from nodes 1-2, 3-4, 5-6 and 7-8 from 12-15 individuals. For the other
10 tissues (seed, stem, leaf, floral organs) each replicate contained material from 7-9 individuals. Traces of DNA were eliminated with TURBO DNA-free® (Ambion). Two µg of RNA were used to make cDNA with the High-Capacity cDNA Archive Kit (Applied Biosystems). Quantitative PCR was performed with SYBR® Green PCR Master Mix (Applied Biosystems) and the Applied
15 Biosystems 7500 real-time PCR system, according to the manufacturer's instructions. Primers used are in Table S2. Three biological replicates were analyzed in each case. Ct values were obtained with the 7500 Systems SDS software 1.3.1.21 (Applied Biosystems). Relative fold expression changes were calculated as described in (Aguilar-Martinez *et al.* 2007. Plant Cell, 19: 458-
20 472). ACTIN8 whose expression levels are constant in all the tissues and conditions analyzed was used as a normalizer. In all the figures the calibrator is the sample with the highest expression.

Isolation of *BRC1*-like genes from wild tomato species

25 Young leaf tissue was collected in N(I), grinded and used to extract genomic DNA with the DNeasy Plant Mini kit (QIAGEN). To amplify the coding regions of *BRC1a* and *BRC1b* inventors carried out nested Pwo polymerase (ROCHE) PCR reactions according to manufacturer's instructions using primers indicated
30 in Table S2.

Phylogenetic analyses

TCP domain-coding DNA sequences were aligned with MUSCLE (v3.7) and default parameters (Edgar 2004. Nucleic Acids Res, 32: 1792-1797). Test of best nucleotide substitution evolutionary model was done with MODELTEST.

- 5 The best fit model (AIC selection) was K81+G+I (w parameter for gamma distribution=0.725). Maximum likelihood (ML) tree reconstruction with the best model and 1000 bootstrap pseudoreplicates was run in PhyML (v3.0 aLRT). The complete phylogenetic pipeline was run in Phylemon web server. Tree was represented with TreeDyn (v198.3).

CLAIMS

1. An isolated polynucleotide, encoding a polypeptide comprising:
 - a. an amino acidic sequence from *Solanum pennellii* which is at least
5 95% identical over the full length of SEQ ID NO: 1 or of SEQ ID
NO: 2, or
 - b. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1
or SEQ ID NO: 2.
- 10 2. An isolated polynucleotide, encoding a polypeptide consisting of:
 - a. an amino acidic sequence from *Solanum pennellii* which is at least
95% identical over the full length of SEQ ID NO: 1 or of SEQ ID
NO: 2, or
 - b. the amino acidic sequence from *Solanum pennellii* SEQ ID NO: 1
15 or SEQ ID NO: 2.
3. The polynucleotide according to claim 2, wherein said polynucleotide is
 - a. SEQ ID NO: 3 encoding SEQ ID NO: 1, or
 - b. SEQ ID NO: 4 encoding SEQ ID NO: 2.
- 20 4. The polynucleotide according to claim 2, wherein said polynucleotide is
the genomic sequence SEQ ID NO: 5.
5. The polynucleotide according to any of claims 1 to 4 operatively linked to
25 any regulatory sequence that controls the expression of said
polynucleotide, preferably the regulatory sequence is SEQ ID NO: 6,
SEQ ID NO: 7 or CaMV35S.
6. An expression product of the polynucleotide according to any of claims 1
30 to 5.

7. An expression vector comprising the polynucleotide according to any of claims 1 to 5.
8. A host cell, having the polynucleotide according to any of claims 2 to 4;
5 having the polynucleotide according to any of claims 2 to 4 operatively linked to any regulatory sequence that controls the expression of said polynucleotide, being preferably said regulatory sequence SEQ ID NO: 6, SEQ ID NO: 7 or CaMV35S; having the expression product of the polynucleotide according to any of claims 2 to 4; or having the
10 expression vector comprising the polynucleotide according to any of claims 2 to 4 or comprising the polynucleotide thereof operatively linked to the regulatory sequence, wherein if the host cell is a plant cell, said plant cell is not a *Solanum pennellii* cell.
9. A use of the polynucleotide according to any of claims 1 to 5, the expression product according to claim 6, the expression vector according to claim 7, or the host cell according to claim 8, to produce a tomato plant or germplasm thereof but not a *Solanum pennellii* plant or germplasm thereof, having fewer and/or shorter axillary shoots than a control plant.
15
10. The use according to claim 9, wherein the produced tomato plant, or germplasm thereof, is classified as *Solanum lycopersicum* species.
20
11. A tomato plant, but not of the species *Solanum pennellii*, having the polynucleotide according to any of claims 2 to 4; having the polynucleotide according to any of claims 2 to 4 operatively linked to any regulatory sequence that controls the expression of said polynucleotide, being preferably said regulatory sequence SEQ ID NO: 6, SEQ ID NO: 7 or CaMV35S; having the expression product of the polynucleotide
25 according to any of claims 2 to 4; having the expression vector comprising the polynucleotide according to any of claims 2 to 4 or comprising the polynucleotide thereof operatively linked to the regulatory
30

sequence; or comprising the host cell according to claim 8, wherein said plant has fewer and/or shorter axillary shoots than a control plant.

5 12. The tomato plant according to claim 11, wherein said tomato plant is a *Solanum lycopersicum* plant.

10 13. The tomato plant according to any of claims 11 or 12, wherein the expression of the polynucleotide is greater than the expression of the homologous native gene *BRC1a*.

14. A germplasm of the tomato plant according to any of claims 11 to 13, preferably the germplasm is a seed or pollen.

15 15. A method for producing tomato plants having fewer and/or shorter axillary shoots than a control plant, comprising:

- a. Transforming at least a tomato plant cell with,
 - i. the polynucleotide according to any of claims 2 to 4, heterologously, or
 - 20 ii. the polynucleotide according to any of claims 2 to 4 operatively linked to any regulatory sequence that controls the expression of said polynucleotide, being preferably said regulatory sequence SEQ ID NO: 6, SEQ ID NO: 7 or CaMV35S, or
 - 25 iii. the expression product of the polynucleotide according to any of claims 2 to 4, or
 - iv. the expression vector comprising the polynucleotide according to any of claims 2 to 4 or comprising the polynucleotide thereof operatively linked to the regulatory sequence, and
- 30 b. growing the plant cell obtained in the step (a) in a suitable medium to produce at least a plant which expresses the heterologous polynucleotide.

16. The method according to claim 9, wherein the tomato plant is a *Solanum lycopersicum* plant.

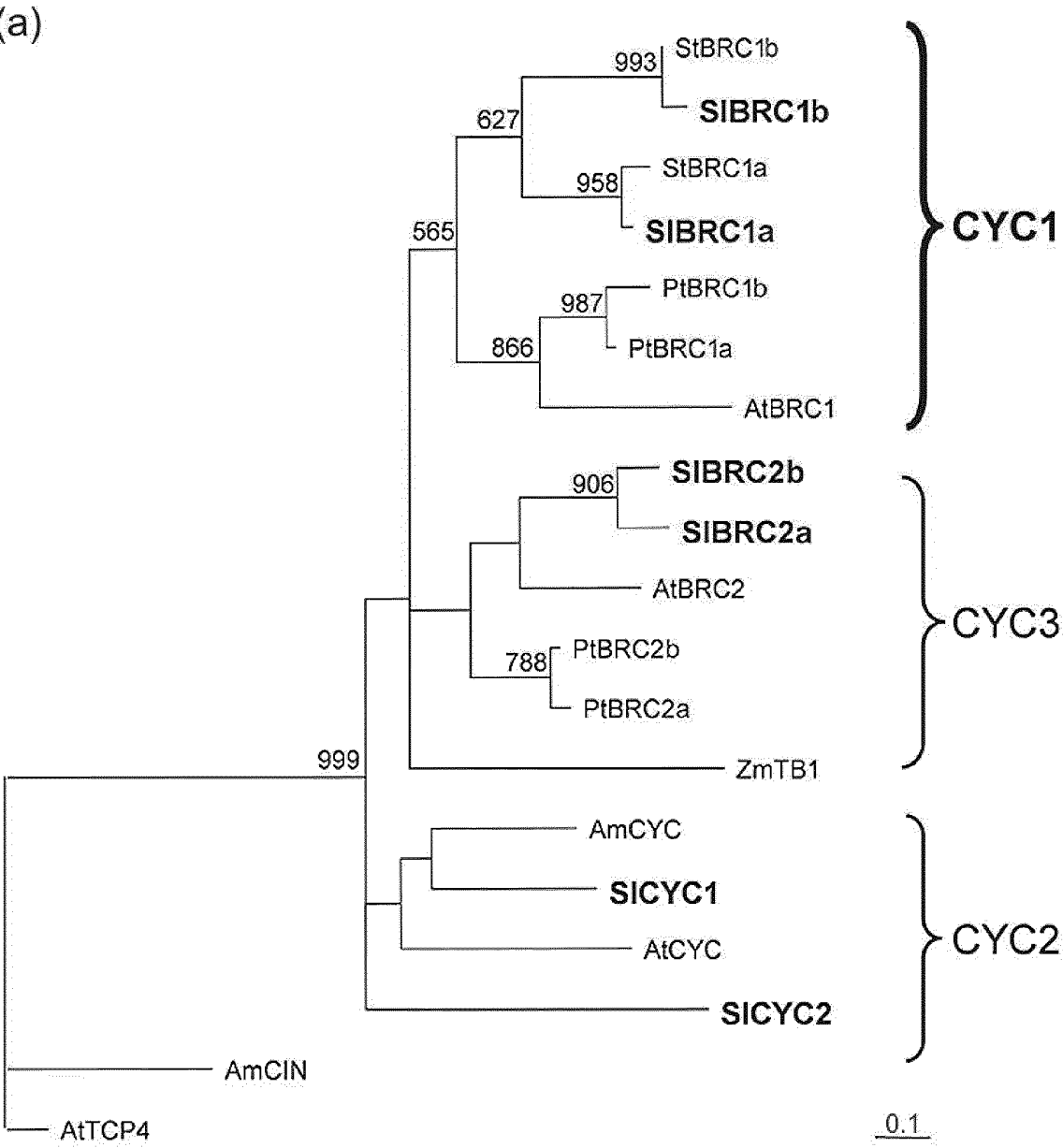


FIG. 1a

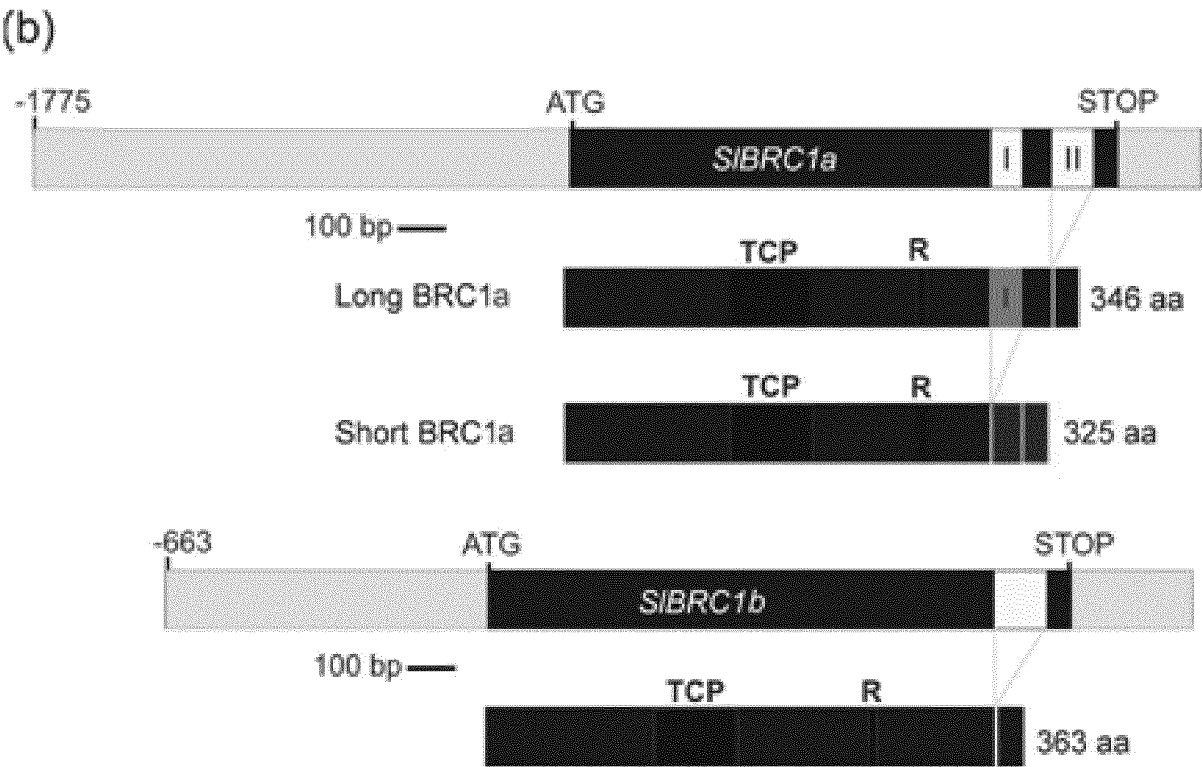


FIG. 1b (Cont.)

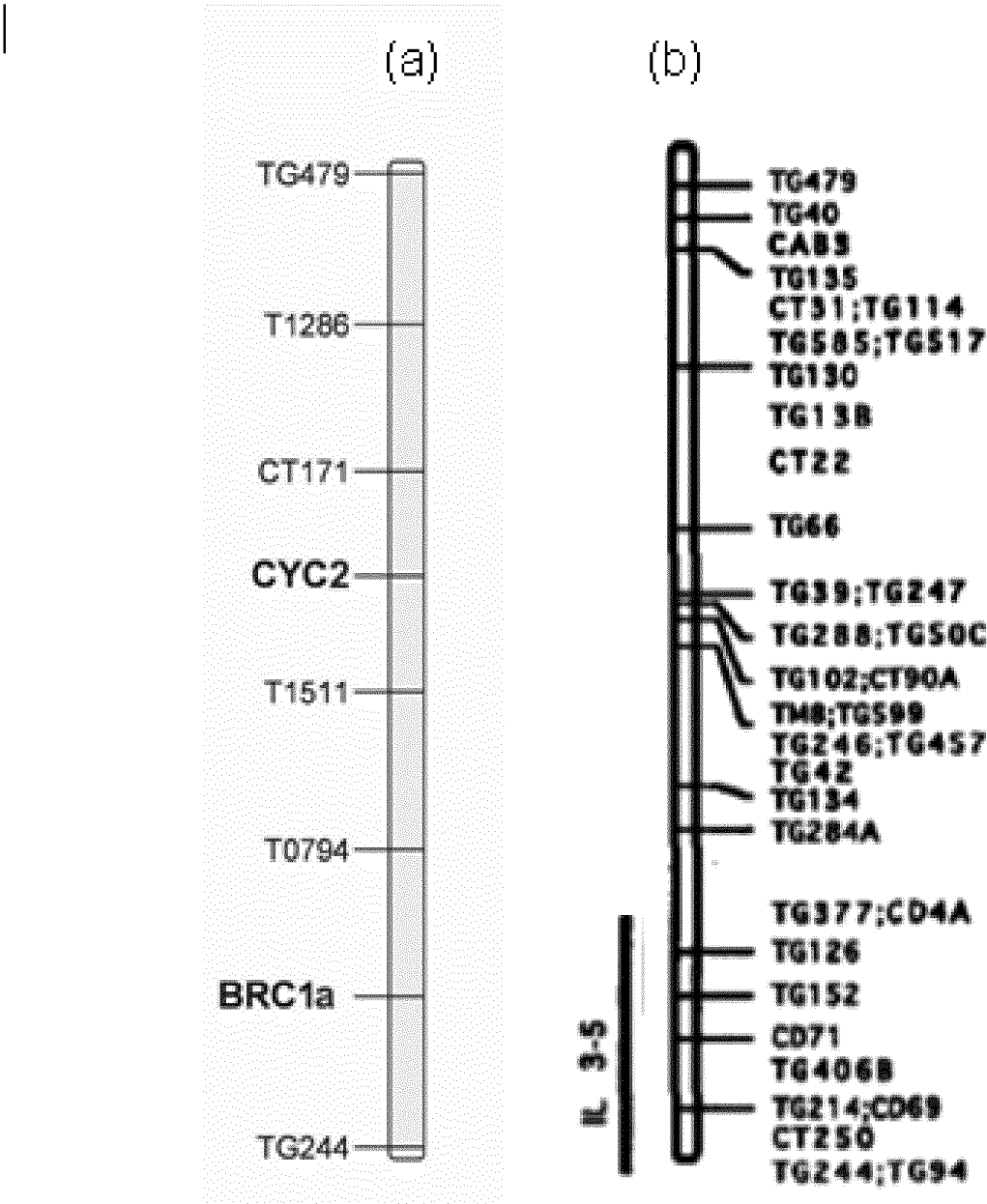


FIG. 2


```

: 1-----11-----21-----31-----41-----51-----
Long BRC1a : IQGYNNNNMMVVDFNLFVDTSNUSPFMFNYHQINTEISQEHQFANFRYSGKLWEA

Jnet      : -----EEEE-----EEE-----
jham      : -----EEEE-----EEE-----

Jnet_25   : ---B---B-BB--B-BB-BB-BB-BB-BB-B-B-B-B-
Jnet_5     : ----B----B-----B-----B-----
Jnet_0     : ----B-----B-----B-----B-----
Jnet_Rel  : 99877788734775254146677776335413234442344442356777665678

: 1-----11-----21-----31-----
Short BRC1a : AHLCSTITKSILKFLKSINLRTSSILGSYGKLN

Jnet      : --HHHHHHHHHHHHH-----HH-----
jham      : --HHHHHHHHHHHHH-----HH-----

Jnet_25   : ---B---B-BBB-B--BB-B--BBB-B--B---
Jnet_5     : ----B---B---B-B-----B-----
Jnet_0     : ----B-----B-----B-----B-----
Jnet_Rel  : 99489999999999998753786421103677899

```

FIG. 3

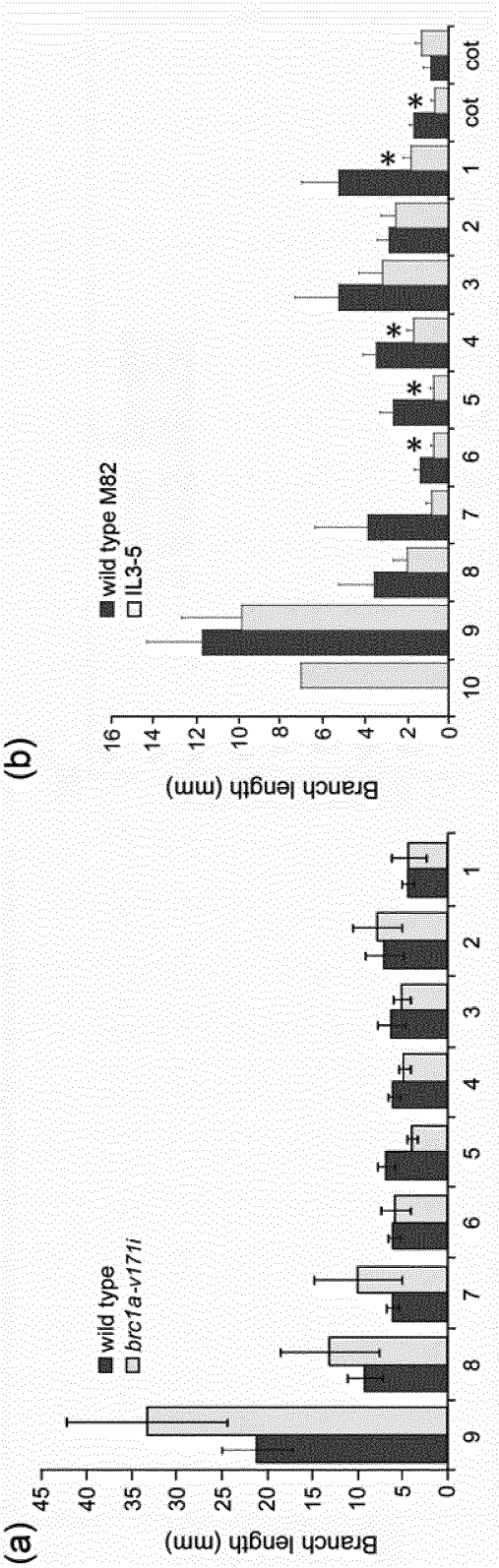


FIG. 4 a-b

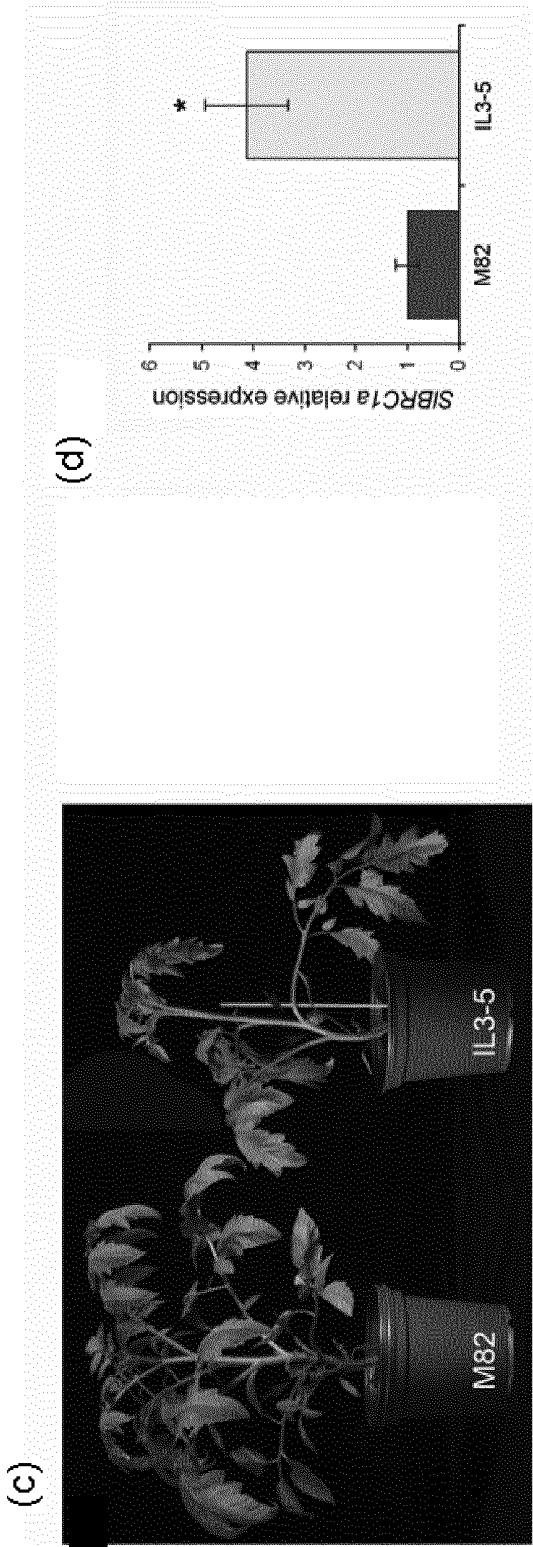


FIG. 4 c-d (Cont.)

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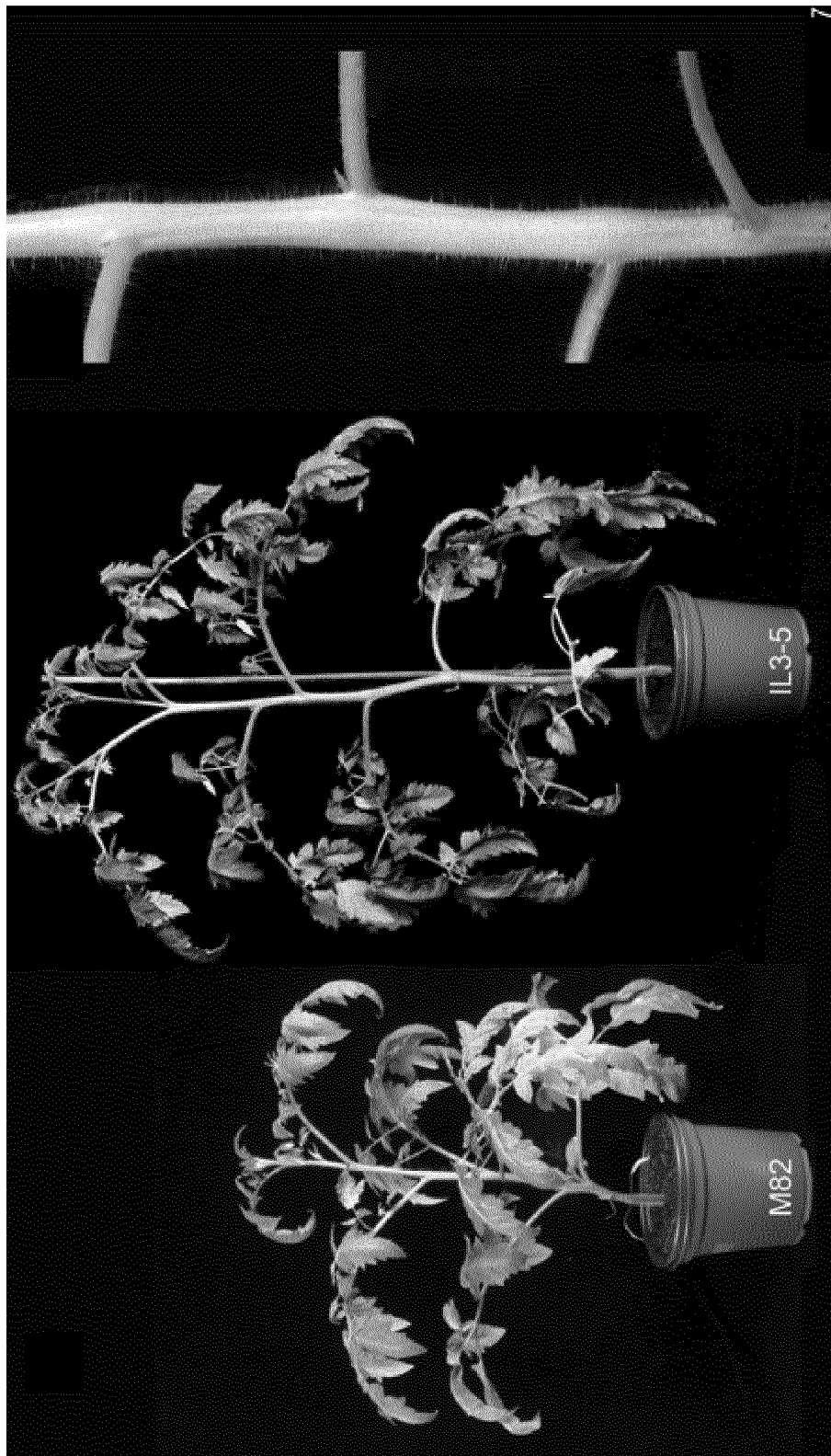


FIG. 5

INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2012/058892

A. CLASSIFICATION OF SUBJECT MATTER

INV. C07K14/415 C12N15/82 C12N5/10 A01H5/00
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C12N A01H C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EP0-Internal, BIOSIS, Sequence Search, PAJ, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	ESHED YUVAL ET AL: "An introgression line population of <i>Lycopersicon pennellii</i> in the cultivated tomato enables the identification and fine mapping of yield-associated QTL", GENETICS, GENETICS SOCIETY OF AMERICA, AUSTIN, TX, US, vol. 141, no. 3, 1 January 1995 (1995-01-01), pages 1147-1162, XP002573291, ISSN: 0016-6731 the whole document page 1161, right-hand column, paragraph 3 -/--	8,11-16



Further documents are listed in the continuation of Box C.



See patent family annex.

* Special categories of cited documents :

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier application or patent but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search

21 June 2012

Date of mailing of the international search report

03/07/2012

Name and mailing address of the ISA/

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Authorized officer

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INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2012/058892

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
	<p>& C.M. RICK: "Second generation Solanum pennellii (syn. L. pennellii) introgression lines", TGRC - TOMATO GENETICS RESOURCE CENTER, 15 April 2002 (2002-04-15), Retrieved from the Internet: URL:http://tgrc.ucdavis.edu/pennellii_ils.aspx> [retrieved on 2011-09-14] -----</p>	
A	<p>WO 2010/081917 A1 (CONSEJO SUPERIOR INVESTIGACION [ES]; MARTIN TRILLO MAR [ES]; RODRIGUEZ) 22 July 2010 (2010-07-22) the whole document -----</p>	1
A	<p>AGUILAR-MARTINEZ JOSE ANTONIO ET AL: "Arabidopsis BRANCHED1 acts as an integrator of branching signals within axillary buds", PLANT CELL, vol. 19, no. 2, February 2007 (2007-02), pages 458-472, XP002659265, ISSN: 1040-4651 the whole document -----</p>	1
A	<p>CUBAS P ET AL: "The TCP domain: a motif found in proteins regulating plant growth and development", PLANT JOURNAL, BLACKWELL SCIENTIFIC PUBLICATIONS, OXFORD, GB, vol. 18, no. 2, 1 January 1999 (1999-01-01), pages 215-222, XP002419545, ISSN: 0960-7412, DOI: 10.1046/J.1365-313X.1999.00444.X the whole document -----</p>	1
A	<p>SCHMITZ G ET AL: "Genetic control of branching in Arabidopsis and tomato.", CURRENT OPINION IN PLANT BIOLOGY FEB 1999 LNKD- PUBMED:10047573, vol. 2, no. 1, February 1999 (1999-02), pages 51-55, XP002659266, ISSN: 1369-5266 the whole document -----</p>	1
T	<p>MARTIN-TRILLO MAR ET AL: "Role of tomato BRANCHED1-like genes in the control of shoot branching", PLANT JOURNAL, vol. 67, no. 4, 21 June 2011 (2011-06-21), pages 701-714, XP002659267, the whole document page 705; figure 2 -----</p>	

INTERNATIONAL SEARCH REPORT

International application No.

PCT/EP2012/058892

Box No. I Nucleotide and/or amino acid sequence(s) (Continuation of item 1.c of the first sheet)

1. With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, the international search was carried out on the basis of:
 - a. (means)
☐ on paper
☒ in electronic form
 - b. (time)
☒ in the international application as filed
☐ together with the international application in electronic form
☐ subsequently to this Authority for the purpose of search
2. ☐ In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.
3. Additional comments:

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No

PCT/EP2012/058892

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 2010081917 A1	22-07-2010	CN 102348801 A	08-02-2012
		EP 2383344 A1	02-11-2011
		ES 2351009 A1	31-01-2011
		US 2012017337 A1	19-01-2012
		WO 2010081917 A1	22-07-2010
